Plasticizers from Lactic Esters and Dibasic Acids

C. E. REHBERG, T. J. DIETZ¹, P. E. MEISS², AND MARION B. DIXON Eastern Regional Research Laboratory, Philadelphia 18, Pa.

Lactic acid can be made economically by the fermentation of agricultural wastes and by-products containing fermentable sugars. As part of a long-range program intended to develop the industrial utilization of lactic acid, many dicarboxylic acid derivatives of lactate esters have been screened to determine their suitability as plasticizers.

Preliminary evaluation of numerous high boiling diesters of lactic acid showed that most of the esters were compatible with polyvinyl chloride and with cellulose acetate. Many of the esters were more efficient than di-2-ethylhexyl phthalate as plasticizers for polyvinyl chloride. The esters that appear to be of most interest as plasticizers for vinyl chloride copolymer are the adipates and sebacates of the butyl and octyl lactates. The direct esterification of alkyl lactates with dicarboxylic acids was also accomplished. In the reaction of adipic acid with butyl lactate, a mixture of esters was produced. These were separated, identified, and evaluated separately as plasticizers for vinyl resins. It was possible to control in large degree the proportions of these various esters in the product. A practical process was developed for the production of a high boiling, substantially neutral, light-colored mixture of esters which was an efficient plasticizer for the vinyl resin and did not require distillation before use. This process was readily adaptable to the use of other lactates and other dicarboxylic acids.

The potential low cost of lactic acid, its ready availability, and the desirable properties of certain derivatives with dicarboxylic acids make these esters of interest to the plasticizer industry.

NTEREST in the utilization of lactic acid in the manufacture of plasticizers has been intensified in recent years by published data showing that lactic acid can be produced at a cost competitive with currently used plasticizer intermediates (3) and that highly efficient plasticizers can be made from lactic acid (7, 8). Recurrent shortages and threats of shortages of key intermediates currently used in plasticizer manufacture have stimulated interest in the use of lactic acid as an extender for scarce materials. Since lactic acid is both an acid and an alcohol, it can be used as an ex-

tender of either the acidic or the alcoholic component of conventional ester type plasticizers.

Previous papers in this series reported on the suitability as plasticizers of esters of lactic acid acylated with alkyl chloroformates (8) and diethylene glycol bischloroformate (7). This paper describes similar esters of adipic, phthalic, maleic, succinic, sebacic, carbonic, and benzenephosphonic acids. These esters have the type formula X[COOCH(CH₂)COOR]₂ where X(COOH)₂ is the dibasic acid used and CH₂CHOHCOOR is the lactate used to make the compound. Most attention was devoted to the adipic and phthalic esters because these acids are available at low cost and are widely used in the plasticizer industry.

¹ Present address, Delaware Research and Development Corp., New Castle, Del.

² Present address, Thiokol Corp., Trenton, N. J.

| | | | | | | | | Prop | erties of Pla | sticized Pol | yvinyl Chlo | ride ^a | ibility with |
|--|--------------------------------------|--|---|---|---|---|--|---|--|--|--|--|--------------------------------|
| Ester Lactate used | Dibasic acid used | 0.01 mm. | Boiling Point 0.1 mm. | ts, ° C | 10 | . d ²⁰ | Viscos- ity, 20°, Cp. | Compatibility b | Tensile strength, lb./sq. inch | Elonga- tion, % | Modulus (100%), lb./sq. inch | Brittle point, ° C. | Cellu- lose Ace- tate |
| Ethyl Propyl Butyl Isobutyl sec-Butyl 2-Ethylhexyl n-Octyl 3.5,5-Trimethylhexyl Allyl 2-Butoxyethyl 2-Hexyloxyethyl 2-(2-Butoxyethyx) 1-Carbethoxyethyl 1-Carbethoxyethyl 1-Carbobutoxyethyl Methyld Ethyl's Butyl 2-Ethylbutyl n-Octyl n-Octyl 2-Ethylhexyl 2-Ethylhexyl 2-Butoxyethyl 2-Butoxyethyl 2-Butoxyethyl 2-Butoxyethyl 2-Butoxyethyl 2-C2-Butoxyethyl | Adipic | 100 105 122 118 106 160 156 154 111 159 165 171 118 135 176 166 166 166 167 173 | 136 136 155 149 138 193 202 188 199 144 192 213 224 200 188 208 145 150 169 187 212 203 191 209 218 | 168 174 194 188 177 233 243 227 239 185 235 268 240 227 186 190 208 228 224 234 225 234 245 245 245 245 245 245 245 245 245 24 | 214 221 240 235 242 280 292 272 288 232 282 | 1 .1075 1 .0865 1 .0544 1 .0479 1 .0574 0 .9960 0 .9981 1 .1085 1 .1085 1 .1085 1 .1077 1 .1478 1 .1072 1 .1074 1 .1094 1 .0878 1 .0878 1 .0878 1 .1094 1 .0878 1 .1094 1 .0878 1 .1094 1 .1092 1 .1092 1 .1092 1 .1092 | 49.65 41.4 53.3 75.6 68.0 53.7 82.4 117.0 42.0 71.9 66.5 526 544 7495 284 284 284 284 284 284 284 284 284 284 | CBB CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC | 3420 3480 3060 3390 3390 3070 3010 3150 3030 3230 3100 3080 3520 3680 3570 3780 3780 3580 3580 3580 3580 3580 3580 3580 35 | 350 340 330 340 360 360 360 325 300 370 280 370 220 290 310 340 325 330 370 220 280 310 280 310 280 310 310 310 310 310 310 310 31 | 1370 1370 1370 1170 1340 1580 1255 1270 11470 1140 1230 1180 1230 1270 2230 1910 2500 2500 240 2300 1910 2020 2110 | -28 -24 -30 -16 -20 -44 -40 -31 -22 -37 -38 -41 -7 -11 -7 -13 9 9 1 -2 -11 -2 -2 -11 -2 -2 -11 -2 -2 -11 -2 -2 -12 -1 | CCCCCHITICCICCCCCHI |
| Allyl Tetrahydrofurfuryl Ethyl Butyl n-Octyl 2-Octyl Methyl Ethyl Butyl n-Octyl And Allyl | Sebacic Succinic | 128 182 128 148 186 172 80 87 107 158 98 | 162 220 160 182 223 208 110 117 139 191 | 203 265 199 222 266 249 146 153 178 231 167 | 257 244 270 190 198 227 281 214 | 1.1693 1.2188 1.0549 1.0200 0.9782 0.9740 1.1925 1.1344 1.0714 1.0042 1.1353 | 298 351 2009 58.3 48.7 63.0 90.9 140.6 54.0 44.7 56.8 48.9 | C C B C C C I C C C I | 3820 3740 3610 3130 2970 3080 2920 3300 3760 3310 3070 2860 | 310 255 240 340 350 350 220 250 310 320 370 | 2010 2780 3060 1290 1190 1360 1230 2550 2080 1410 1440 1230 | -14 -4 19 -28 -31 -40 -38 -21 -29 -23 | 00000011000010000100 |
| Ethyl Butyl Isobutyl 2-Ethylhexyl Allyl Butyl Allyl Di-2-ethylhexyl phthalate (Control) | Maleic Benzene- phosphonic Carbonic | 60 105 97 150 100 130 72 120 | 89 136 127 (0.03 mm.) 131 (0.02 mm.) 101 153 | 124 174 166 169 135 192 | 170 221 212 | 1.1244 1.0852 1.0847 1.0134 1.1562 1.1082 | 22.9 74.3 189.5 110.4 | I C C C C B C B | 3590 / >3240 3130 3450 / 3530 3130 3800 3070 | 310/ >330 365 340/ 340 350 310 290 | 2230 / 1670 1620 / 1610 / 2140 1490 | -23 -7' -19 -12 -36' -22 -11 | CI CC CI |

- a A 95:5 copolymer of vinyl chloride and vinyl acetate containing 35% plasticizer.

 b C = Compatible; I = incompatible; CI = borderline; B = compatible when milled but bled on aging.

 d High acetyl cellulose acetate containing 20% plasticizer.

 d Melting point, 50-55° C.; n_D and d on supercooled liquid.
- Melting point, 55-60° C.; n_D and d on supercooled liquid.
- f Properties determined with undistilled material.

PREPARATION AND PROPERTIES OF ESTERS

Table I shows the esters studied and the properties determined. Additional data on the preparation and properties of these esters have been published elsewhere (5, 6).

The boiling points shown in Table I were read from the Cox charts previously published (5, 6). Most of the esters had boiling points higher than that of di-2-ethylhexyl phthalate. Comparison of the vapor pressure curves of the esters in Table I with those of the corresponding esters not containing lactate radicals for example, bis(butyl lactate), adipate with dibutyl adipateshows that the vapor pressure of the esters containing lactate radicals is only 3 to 5% of that of the corresponding simple esters. Additional data on densities and viscosities are given in references (5) and (6).

Evaluation as Plasticizers. Most attention was devoted to polyvinyl chloride because this resin accounts for the largest volume of plasticizers, and its use is still expanding. The compounding and testing procedures have been described previously (7), and the results are summarized in Table I. As judged by modulus and brittle point, about one fourth of the esters are more efficient plasticizers for the 95% vinyl chloride copolymer than is di-2-ethylhexyl phthalate. Nearly all milled well and appeared compatible when tested, but several samples showed "bleeding" or spewing after being kept several months. Possibly some of these, which otherwise appear attractive as plasticizers, could be used in blends, thus improving their limited compatibility.

The presence of rings in the molecular structure of the esters markedly raised the brittle points of the plasticized vinyl resin, as is illustrated by the tetrahydrofurfuryl and the phthalate esters. A similar effect is produced by lactyllactate radicals, as in ethyl and butyl lactyllactate adipates-1-carbethoxyethyl and 1carbobutoxyethyl, lactate adipate, respectively. The esters that appear to be of most interest as plasticizers for vinyl resin are the adipates and sebacates of the butyl and the octyl lactates. They combine good compatibility with high efficiency and exceptional permanence.

Compat-

Compatibility of the esters with high acetyl cellulose acetate was determined by casting plasticized films from solution in acetone. As may be seen in Table I, most of the esters were compatible at 20% concentration. In many instances the presence of the lactate radicals in the plasticizers has greatly improved the compatibility with cellulose acetate. At the same time, of course, volatility is much reduced, thus providing a unique group of very high boiling compatible plasticizers for cellulose acetate.

DIRECT ESTERIFICATION OF ALKYL LACTATES WITH DICARBOXYLIC ACIDS

In the preparation of small amounts of these compounds (listed in Table I) for screening purposes the dibasic acid chlorides were used in most cases as a matter of convenience (5, 6). This method was obviously impractical for commercial production of the compounds, and so a study was made of the reaction of dicarboxylic acids with alkyl lactates.

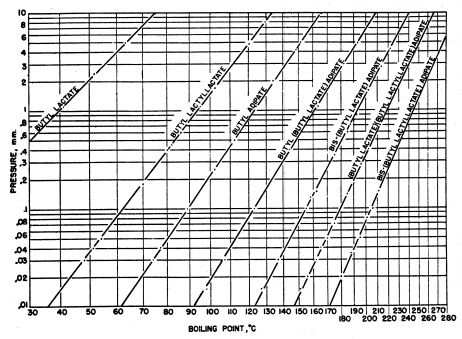


Figure 1. Relationship of Boiling Points to Pressure for Various Esters

On the basis of desirability, cost, ease of preparation, and availability of intermediates, butyl lactate adipate appeared the most attractive of the diesters screened. Hence, in the present work, most attention was devoted to the esterification of butyl lactate with adipic acid. A few experiments were run in which butyl or capryl lactate was esterified with sebacic, maleic, or phthalic acid or anhydride.

Since a lactate ester is also an alcohol, it is capable of undergoing self-alcoholysis when heated in the presence of strong acidic or basic catalysts (1). This results in the formation of linear polyesters:

2CH₃CHOHCOOR
$$\longrightarrow$$
 CH₃CHOHCOOCH(CH₃)COOR + ROH
 n CH₃CHOHCOOR \longrightarrow HO[CH(CH₃)COO]_n - R + $(n-1)$ ROH

Thus in the esterification of adipic acid with butyl lactate, for instance, the alcoholic hydroxyl groups that react with the acid may be furnished by butanol, butyl lactate, butyl lactyllactate, or some higher polylactic ester. The situation is further complicated by the fact that dicarboxylic acids may form unsymmetrical esters with any two of the alcoholic compounds present.

Fractional distillation of the mixture of esters formed from adipic acid and butyl lactate resulted in the isolation of the principal components as relatively pure compounds:

```
COOCH(CH<sub>2</sub>)COOCh(CH<sub>2</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(CH<sub>3</sub>)COOCh(
```

COOCH(CH₃)COOCH(CH₃)COOC₄H₉ (CH₂)₄ COOCH(CH₃)COOCH(CH₃)COOC₄H₉ Bis(butyl lactyllactate)adipate

These esters are shown in Table II with some properties of the crude mixtures of esters. The symmetrical esters, bis(butyl lactate) and bis(butyl lactyllactate) adipate were also prepared in pure form by use of adipyl chloride. The properties shown for these in Table II were determined on these pure samples (δ) . The unsymmetrical esters were isolated from the mixtures of esters obtained by use of adipic acid and hence were somewhat less pure.

Figure 1 is a Cox chart in which the boiling points [1/(t+273)] are plotted versus the pressure $(\log P)$, t being in $^{\circ}$ C. and P in mm. This figure was a most useful guide in the distillation of the mixtures of esters obtained in the various experiments.

Study of Esterification Reaction.

A brief study was made of the effect

of several variables on the amount and kind of products formed in the acid-catalyzed esterification of butyl lactate with adipic acid. In these experiments the reactants were refluxed in a still having a 24-inch column packed with glass helices and having a water trap above the packing. Heating was begun at atmospheric pressure, steady reflux was maintained, water was removed as formed (butanol formed in the reaction served as an azeotropic agent to remove the water), and the reaction temperature was kept at or below 150° C. by suitably reducing the pressure from time to time as the reaction progressed. When reaction appeared complete, the catalyst was neutralized (preferably with a slight excess of powdered anhydrous sodium acetate), and butyl alcohol, lactate, lactyllactate, and adipate were removed successively by distillation at reduced pressure through a $1^{1/2} \times 24$ inch Vigreux column. The residue, after washing with dilute alkali and water, drying, and filtering, was suitable for use as a plasticizer.

The composition of the residue was estimated by distilling all or an aliquot portion in a high vacuum alembic still (4) and analyzing the fractions by boiling point and refractive index. Table III summarizes the results obtained in some selected typical experiments.

Experiments 1 and 2 (Table III) show that increasing the amount of catalyst markedly reduces the time required for reaction. The use of an equivalent amount of toluenesulfonic acid instead of sulfuric acid in several runs, including Expt. 10, made little difference in reaction time or color and composition of the product.

Comparison of the results of Expt. 2 with 3 or 8 with 9 shows that by increasing the proportion of butyl lactate in the reaction mixture above the theoretical amount, relatively more of the products boiling below bis(butyl lactate) adipate and less of the higher boiling products are produced.

The production of butyl adipate in the reaction mixture was undesirable because its high volatility necessitated its removal from the product before the latter could be used as a plasticizer. On the assumption that the composition of the reaction mixture approached a point of dynamic equilibrium, it was expected that addition of butyl adipate to the reaction mixture would reduce or prevent the formation of this compound during the reaction.

In Expts. 4 to 7, an increasing proportion of butyl adipate was

TABLE II. PROPERTIES OF ESTERS FORMED FROM ALKYL LACTATES AND DIBASIC ACIDS

| TABLE II. Pro | PERTIE | s of E | STERS | FORMED F | HOM ALE | ID DAOLL. | | | | | |
|--|---|--|--|---|---|---|---|--|---|---|--|
| Compound Butyl lactyllactate Butyl adipate Butyl (butyl lactate) adipate (Butyl lactate); adipate (Butyl lactate); adipate (Butyl lactate); adipate (Butyl lactate); adipate Crude butyl lactate adipate Capryl (capryl lactate) adipate (Capryl lactate); adipate Crude capryl lactate adipate Crude capryl lactate adipate Crude 2-ethylhexyl lactate); adipate Crude 2-ethylhexyl lactate adipate Capryl (capryl lactate); sebacate Crude capryl lactate); sebacate Crude capryl lactate; sebacate | | SOF E 12 Points 0.4 mm. 80 109 145 177 202 227 184 211 216 207 232 125 158 | | n20 1.4328 1.4353 1.4396 1.4432 1.4461 1.4470 1.4418 1.4432 1.4446 1.4438 1.4494 1.4494 1.4494 1.4494 1.4495 1.4470 1.4470 1.4470 | d20 1.0631 0.9615 1.0175 1.0543 1.0922 1.1026 1.0458 0.9515 0.9955 0.9955 0.9980 0.9442 0.9740 0.9440 1.0852 | Mol. Ref Calcd. 53.21 70.16 85.67 101.18 116.68 132.19 101.18 122.61 138.12 138.12 138.12 138.12 141.09 156.59 75.97 91.48 | Found 53.34 70.17 85.50 101.24 115.96 132.42 101.77 123.37 137.83 141.58 138.17 140.95 166.74 160.99 76.30 92.01 | Sapon. Calcd 110.1 100.6 94.9 91.1 100.6 147.5 128.7 128.7 128.7 128.7 128.7 128.7 128.7 128.3 142.2 142.2 142.3 | Equiv. Found 108.3 99.7 94.2 91.9 104.7 152.7 125.6 134.2 128.4 135.2 163.3 141.1 152.4 99.8 93.8 | Viscosit 20° C. 16.20 15.55 41.37 177.1 283.5 31.74 21.70 82.4 68.0 45.6 90.9 60.6 22.66 74.3 | 40° C. 6.69 7.80 16.56 46.68 67.88 14.09 10.20 23.55 25.6 19.50 35.4 26.5 19.36 |
| Capryl (capryl lactate) sebacate (Capryl lactate); sebacate Crude capryl lactate esbacate Butyl (butyl lactate) maleate (Butyl lactate); maleate Crude butyl lactate maleate Butyl (butyl lactate) phthalate (Butyl lactate); phthalate Crude butyl lactate phthalate Reaction product was topped to: \$\begin{array}{c} a 110 \\ b \\ 100 \\ c \\ c \\ 175 \\ c \\ c \\ c \\ 175 \\ c \\ c \\ c \\ 175 \\ c \\ c \\ c \\ c \\ 175 \\ c \\ | 90 123 124 155 C. (vapor C. (vapor C. (pot te | 232 125 158 163 191 temp.) s temp.) s | 280 169 201 211 237 at 0.1 mr at 0.2 mr 0.07 mm | 1.4475 1.4470 1.4470 | 0.9740 0.9440 1.0524 1.0852 1.0727 1.0842 1.1094 1.093 ipate-free), ctyllactate- tyllactate- tyllactate- tyllactate- | 156.59 75.97 91.48 91.47 91.30 106.81 106.81 washed, dri- free), washed | 160.99 76.30 92.01 92.74 92.84 108.56 110.80 ed, and filted | 142.2 100.1 93.1 93.1 116.8 105.6 105.6 ered. | 152.4 99.8 93.8 95.5 115.2 105.6 109.2 | 66.6 22.66 74.3 | 26.5 9.65 19.36 |

put into the reaction and, as expected, the amount formed decreased until, in Expt. 7, less butyl adipate was recovered in the product than had been put into the reaction mixture. It is evident, then, that butyl adipate can be recycled in the process so that none remains to be disposed of as a by-product. The same applies to any butyl lactate or lactyllactate recovered from the product.

In all experiments, small amounts of butyl lactyllactate adipates were recovered, and a distillation residue was obtained which probably consisted of higher polylactic esters. Because these polymeric esters were relatively less efficient as plasticizers than the monomeric lactic esters, their production in the esterification reaction was undesirable, and ways to prevent their formation were sought. In comparing Expt. 3 with 1 and 2, it was noted that increasing the proportion of butyl lactate used decreased the amount of high boiling polylactic esters and distillation residue, but only at the price of greatly increasing the amount of lactate and lactyllactate recovered. The unsymmetrical ester, butyl (butyl lactate) adipate, is a highly efficient plasticizer and is sufficiently high boiling for most applications, particularly when used in admixture with the higher boiling lactate adipates. Formation of this ester is favored and at the same time formation of polylactic esters is repressed by addition of butanol to the reaction mixture, as shown by the results of Expts. 8 and 9.

Since lactic acid is a commercially available raw material, it appeared desirable to combine the two esterifications, (a) lactic acid to butyl lactate and (b) butyl lactate to butyl lactate adipate, into a single step. This was done in Expt. 11, and it appeared that substantially the same result was obtained as when butyl lactate was first prepared separately.

According to a recent appraisal (3), the methanol vapor process (2) of purifying lactic acid is probably the most economical way to make high grade lactic acid. Hence, methyl lactate, an intermediate in this process, appeared to be potentially cheaper than lactic acid or butyl lactate. Expt. 12 was run to determine the feasibility of combining the two steps, (a) conversion of methyl lactate to butyl lactate and (b) esterification of butyl lactate with adipic acid, into a single step. Methanol was distilled from the reaction mixture as it was formed, after which water was removed as usual. The product was not significantly different from that prepared from butyl lactate or lactic acid.

Although it seemed evident that the general conclusions derived from the experiments in which adipic acid and butyl lactat were used would be valid for the use of other dicarboxylic acids and other alkyl lactates, a few experiments were run to verify this expectation. 2-Ethylhexyl lactate was esterified with adipic acid, capryl lactate was esterified with adipic and sebacic acids, and butyl lactate was esterified with maleic and phthalic anhydrides. In addition, an experiment analogous to 12, Table III, was run in which sebacic acid was reacted with ethyl lactate and capryl alcohol. No significant difference was noted in the reactivity of the various acids and anhydrides, but it was apparent that capryl alcohol and capryl lactate were less reactive in reactions involving the alcohol radical than the butyl compounds. The esters produced in these experiments are included in Table II.

The color of the reaction products depended to a great extent

| | | , e e 6 | Таві | æ II | I. Ester | rificatio | ON OF | BUTYL I | | | IPIC AC | ID pic Acid Use (Bu. lac- | d ——— | |
|--------------------------|---|---------------------|--|--|--|--|--|--|---|--|--|--|--|--|
| Expt. | Reac | etants, M Adipic | Bu. adi- | —— Bu. | Catalysts, Grams | Reac- tion Time, Hours | Bu. | Bu. lactyl- lactate | Bu. adipate | Bu. (Bu. lactate) adipate | Bis(Bu. lactate) adipate | tate) (Bu. lactyl- lactate) adipate | Bis(Bu. lactyl- lactate) adipate | Total plasti- ciser ^b , moles |
| No. 1 2 3 4 5 6 7 8 9 10 | Bu. Lactate 2 2 4 8 6 4 2 8 10 10 8 (Lactic acid) 8 (Me. lactate | | pate 0 0 1 1 1 2 2 2 | alc. 0 0 0 0 0 0 0 4 4 0 10 | H ₁ 8O ₄ 0.2 1.0 1.0 4.0 3.0 2.0 1.0 4.0 4.0 4.0 4.0 4.0 | 12 5 1.5 6 4 4 3 3 4 4 6 | 0.19 0.17 1.45 0.20 0.11 0.16 0.17 0.63 0.93 0.28 0.28 | 0.10 0.52 0.08 0.05 0.07 0.08 0.30 0.25 0.16 0.08 | 0.23 0.23 0.33 0.29 0.05 0.07 -0.30 0.20 0.38 0.02 0.21 | 0.22 0.28 0.43 0.34 0.43 0.38 0.64 0.51 0.51 0.50 0.42 | 0.21 0.27 0.22 0.20 0.28 0.27 0.23 0.15 0.14 0.31 0.19 | 0.14 0.07 0.06 0.18 0.13 0.11 0.20 0.03 0.03 0.12 0.09 0.11 | 0.07 0.04 0.01 0.04 0.04 0.05 0.01 0.00 0.05 | 0.64 0.66 0.72 0.76 0.88 0.80 1.12 0.70 0.68 0.98 0.98 |

Net amount in excess of that put in.
Total number of moles of ester distilled above butyl adipate.
Toluenesulfonic acid.

ΓABLE IV. PROPERTIES OF PLASTICIZED POLYVINYL CHLORIDE-ACETATE^a

| Plasticizer | Tensile Strength, Lb./Sq. Inch | Ultimate Elonga- tion, | Modulus, (100% Elong.) Lb./ Sq. Inch | Brittle Point, |
|--|---|------------------------------|---|-------------------|
| | | , ,, | | |
| 2-Ethylhexyl phthalate (con- | | | | |
| _ trol) | 3070 | 290 | 1500 | -32 |
| Butyl adipate | >2200 | >390 | 700 | -68 |
| Butyl (bu. lactate) adipate | 2620 | 320 | 920 | $-45 \\ -30$ |
| (Butyl lactate), adipate | 3060 | 330 | 1170 | 80 |
| (Butyl lactate) (butyl lactyl- | 2000 | 320 | 1565 | -17 |
| lactate) adipate | 3600 | 320 270 | 1910 | -17 |
| (Butyl lactyllactate), adipate | 3580 | 270 | 1910 | -13 |
| Mixed adipate esters, dis- | 3110 | 330 | 1120 | -28 |
| | 3110 | 990 | 1120 | -20 |
| Mixed adipate esters, undis- tilled | 3145 | 315 | 1030 | -35 |
| Capryl adipate | 2600 | 270 | 1140 | -66 |
| Capryl (capryl lactate) adi- | 2000 | 210 | 1110 | 00 |
| Date | 3110 | 270 | 1210 | -44 |
| (Capryl lactate); adipate | 3070 | 360 | 1270 | -40 |
| (Capryl lactate); adipate, | 00.0 | 000 | | |
| crude d | 3350 | 250 | 1460 | -36 |
| (2-Ethylhexyl lactate); adi- | 0000 | | | |
| pate | 3020 | 260 | 1255 | -44 |
| (2-Ethylhexyl lactate) adi- | | | | |
| pate, crude | 2970 | 360 | 1530 | -26 |
| Caprylsebacate | 2650 | 240 | 1190 | -62 |
| Capryl (capryl lactate) seba- | | | | |
| cate | 2830 | 340 | 1400 | -49 |
| (Capryllactate); sebacate | 2920 | 350 | 1230 | -38 |
| (Capryl lactate): sebacate, | | | | |
| crude * | 2920 | 340 | 1460 | -43 |
| Butyl (butyl lactate) maleate | >3040 | >380 | 1230 | -35 |
| (Butyllactate): maleate | >3240 | >330 | 1670 | -19 |
| (Butyl lactate): maleate, crude | | | | |
| | 3620 | 350 | 1650 | -17 |
| Butyl phthalate | 2670 | 340 | 980 | -30 |
| Butyl (butyl lactate)phthal- | 0.440 | 0.50 | 1700 | 10 |
| ate | 3440 | 350 | 1580 | - 10 |
| (Butyl lactate): phthalate | 3580 | 310 | 1800 | 1 |

^a A 95:5 copolymer of vinyl chloride and vinyl acetate containing 35%

A 90:5 copolymer of vary
b All distillable material boiling above butyl adipate.
c Crude butyl lactate adipate topped to 110° C. (vapor temp.) at 0.1 mm. pressure (butyl adipate-free), washed, dried, and filtered.
Crude esters topped to:
d 100° C. (vapor) at 0.22 mm., washed, dried, and filtered.
175° C. (pot) at 0.07 mm., washed, dried, and filtered.
f 100° C. (pot) at 0.1 mm., washed, dried, and filtered.

on the purity of the materials used. When c.p. grade reagents were used, the sebacates were faintly yellow; the adipates were pale yellow to light brown; the maleates were a clear, medium brown; and the phthalates were a darker, reddish brown.

Decolorizing carbon was useful in removing some of the color but was found to be much more effective if added to the reaction mixture at the beginning of the esterification and filtered out after all processing had been completed. The use of 1% carbon by weight thus yielded clear, almost colorless products. Several brands of activated carbon were used, but no great differences in effectiveness were noted.

Free acidity in the reaction mixtures was followed by withdrawal and titration of samples. It could be reduced to around 1% (as acetic acid) by continuation of the esterification reaction until this value was reached. Further reduction was so slow as to be impractical. Neutralization of the catalyst with sodium acetate produced acetic acid. This was removed in stripping volatile materials from the product, thus further reducing free acidity.

A single wash with 1% sodium carbonate solution was sufficient to reduce the free acidity to around 0.02%, a value acceptable for most purposes. It was easy to wash the crude esters with the alkali if the initial acidity was not above 1 to 2%. However, if the acidity was as high as 3%, the wash was made tedious by the formation of emulsions.

Evaluation of Esters Obtained by Direct Esterification. Each

of the esters isolated from the crude reaction products as well as the undistilled mixtures of esters was evaluated as a plasticizer for a commercial copolymer of 95% vinyl chloride and 5% vinyl acetate, as previously described (7). In each series of esters (Table IV) each additional lactic acid group introduced into the ester increased the tensile strength, modulus, and brittle point of the resin plasticized with it. Of course, the boiling points were increased at the same time, and this is of especial importance in the adipates and maleates.

In the butyl lactate adipate series of esters, it is apparent that dibutyl adipate is much too volatile, although extremely efficient as a plasticizer. Inclusion of one lactate group greatly reduces the volatility (boiling point about equal to that of hexyl phthalate) while retaining exceptional efficiency. The two lactate groups in bis(butyl lactate) adipate raise the boiling point further and produce a plasticizer equivalent to 2-ethylhexyl phthalate in volatility and low temperature performance and somewhat superior in producing compositions of low modulus. The esters of lactvllactic acid have extremely low volatility but are less efficient than those of monomeric lactic acid.

The same general trends are apparent in the other series of esters, the principal exception being that, with the exception of the maleates, the simple esters containing no lactic acid are sufficiently high boiling to permit them to be left in the undistilled plasticizer mixtures.

In the sebacate series, capryl sebacate is a highly efficient plasticizer but is of borderline compatibility. This defect is reduced or eliminated by the inclusion of lactate groups in the molecule, though at the expense of some loss in efficiency and low temperature flexibility.

Any commercial utilization of these products would probably be based on the use of the undistilled mixture of esters produced in the esterification reaction, after removal of the more volatile components. Hence, it was gratifying to find that these undistilled mixtures were as efficient as the distilled products and that the adipate and sebacate mixtures were equal or superior to 2ethylhexyl phthalate in efficiency and permanence.

ACKNOWLEDGMENT

The authors are indebted to W. E. Palm and H. C. Fromuth for the physical testing of the plasticized vinyl resins.

LITERATURE CITED

- (1) Filachione, E. M., Costello, E. J., Dietz, T. J., and Fisher, C. H., U. S. Department of Agriculture, Bureau of Agricultural and Industrial Chemistry: AIC-295, February 1951. (Processed).
- (2) Filachione, E. M., and Fisher, C. H., Ind. Eng. CHEM., 38, 228 (1946).
- (3) Needle, H. C., and Aries, R. S., Sugar, 44, No. 12, 32 (1949).
- (4) Ratchford, W. P., and Rehberg, C. E., Anal. Chem., 21, 1417 (1949).
- (5) Rehberg, C. E., and Dixon, M. B., J. Am. Chem. Soc., 72, 5757 (1950).
- 6) Ibid., 74, 707 (1952).
- Rehberg, C. E., Dixon, M. B., Dietz, T. J., and Fisher, C. H., IND. ENG. CHEM., 42, 1409 (1950).
- Rehberg, C. E., Dixon, M. B., Dietz, T. J., and Meiss, P. E., Ibid., 42, 2374 (1950).

RECEIVED for review September 20, 1951. ACCEPTED May 3, 1952. Most of the material in this paper was presented before the Division of Paint, Varnish, and Plastics Chemistry at the 113th, 116th, and 117th National Meetings of the American Chemical Society. One of the Laboratories of the Bureau of Agricultural and Industrial Chemistry, Agricultural Research Administration, United States Department of Agriculture